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- Process for the conversion of oldfins having from 4 to 12 carbon stams to propylane.
- 5) A process for the conversion of olefine having from 4 to 12 carbon atoms into propylene, characterized in that said olefins are brought into contact with a zeolitic compound (optionally in admixture with a binder), selected from silicalities, boralities, chromosilicates and those zeolities ZSM5 and ZSM11 in which SiO₂/A₂O₃ (by moles) is > 350, at a space velocity of from 5 to 200 kg/h of olefins per kg of pure zeolitic compound (binder excluded) and at a temperature of from 400 to 600°C, said zeolitic compound being used as such or in a modified from.



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PROCESS FOR THE CONVERSION OF OLEFINS HAVING FROM 4 TO 12 CARBON ATOMS TO PROPYLENE

The invention concerns a process for the conversion of olefins having from 4 to 12 carbon atoms into propylene. Nowadays huge amounts of olefine cuts, from C₄ to C₁₂, linear or branched, are available throughout the world and they are widely employed for different purposes, such as described for instance in Italian patent publications 24152 A/82, 24550 A/82 and 19292 A/83. Sometimes, however, because of contingent reasons, even outside the chemical field, like for instance transportation difficulties, it would be better to have still further possibilities of use. A promising use of said olefins would be their conversion into propylene and/or ethylene.

However, endurance tests have shown that the excellent initial behavior of some catalysts, like for instance ZSM5 and ZSM11, disappears after some time; after a few weeks conversion and selectivity drop to poor levels.

An object of the invention is to provide a process for catalytically converting olefin cuts $\mathbf{C_4} - \mathbf{C_{12}}$ with high propylene (and optionally also ethylene) yields and with a long life time of the catalyst before substitution or regeneration.

In its most general form, the invention concerns a process for the conversion of olefins having from 4 to 12 carbon atoms into propylene which is characterized in that said olefins are brought into contact with a zeolitic compound (optionally in admixture with a binder) selected from silicalites, boralites, chromosilicates and those zeolites ZSM5 and ZSM11 in which SiO₂/Al₂O₃ (by moles) is \$\grace\$350, at a space velocity of from 5 to 200 kg/h of olefins parkg of pure zeolitic compound, binder excluded, and at a temperature of from 400 to 600°C, said zeolitic compound being used as such or in a modified form.

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The behavior of silicalites depends on the conversion pressure; for instance, if the silicalite is silicalite-1 and if the pressure is substantially atmospheric, the space velocity must be lower than 50 h^{-1} ; if said compound is silicalite-1 and if the pressure is from 1.5 to 7.5 absolute atmospheres the space velocity must be in general above 50 h^{-1} .

Best results are obtained when said catalytic silicalite-1 is activated in the conversion reaction of the olefins into propylene, under said operative conditions. This initial (activating) run takes at least 8 and preferably 12 hours; the silicalite-1 is used as such or in a modified form and the modifying element is selected from Cr, Mg, Ca, Sr and Ba.

The modifying element can be incorporated into the catalyst by means of ion exchange or by any other method, for instance impregnation or co-precipitation during the synthesis of the zeolitic compound.

According to a particularly advantageous way for the preparation of a non-modified silicalite, the raw product coming from the zeolite synthesis is dried, for instance at 120°C, calcined (e.g. at 540°C for some hours), in order to remove all the residual organic templating agent, and then exchanged with an aqueous solution of HCl, NH₄Cl, NH₄NO₃ or an equivalent H⁺ or NH₄ source. When an ammonium compound is used, it is necessary to heat, e.g. at 400°C, in order to obtain the acid form of the silicate. A survey of techniques alternative to ion exchange is given e.g. in U.S. patents 3,140,249; 3,140,251; 3,140,253 and in European patent publications 30796, 36707, 37168, 40463, 68754.

The zeolitic material, after calcination and conversion into the acid form, shows a long endurance and a very high catalytic activity. These zeolitic compounds can be used as catalysts (as such or in a modified form) optionally in admixture with suitable amounts of binders, for instance SiO₂ or Al₂O₃. A list of other binders can be found

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e.g. in European patent publication 36707. The regeneration can be carried out in air for some hours, at 400-600°C. A steam regeneration is described in European patent publication 36704 and according to a further and very successful method, the catalyst can be regenerated by a hydrogen treatment.

As to the initial activation of the catalyst, some methods are described in European patent publications 34444 and 35830; in general it is advisable to activate the catalyst for some hours in air, at 450-750°C (preferably 540-700°C). Furthermore the conversion itself (of ole-fins to propylene) has an activating effect on the zeolitic catalyst. In other words, catalyst and reaction affect each other in a mutual, reciprocal and beneficial activating reaction.

Any process for the conversion of more or less heavy olefinic cuts into propylene will be indicated hereafter, as a "post-pyrolysis" process. When the feed of a postpyrolysis process is a mixture of olefins having 4 C atoms, there is a considerable problem to be solved, because the C4 cuts always contain substantial amounts of paraffins, in general, also having 4 C atoms, which paraffins pass the zeolitic bed without taking part in (at 400-600°C) any reaction. Furthermore a small amount of C4 paraffins is produced by the post-pyrolysis process itself. The conversion to C3H6 could be enhanced by a recycle of the nonreacted C_4^- olefins or of the C_4^- olefins formed during the reaction. In such a case, however, an increasing accumulation of n-butane and of isobutane would take place. This drawback could be avoided by a separation of paraffins feeding the reactor but such a from olefins before separation is rather difficult. Butenes and isobutane cannot be isolated by a simple distillation and it is usually necessary to carry out an extractive distillation (a complicated technique), which is particularly burdensome for the ${\rm C_4}^-$ cuts coming from catalytic cracking, where butane and isobutane may account for even 50% of the whole. The problem, however, can be solved in a surprisingly easy way

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by the process according to the invention, when employing an integrated oligomerization step. In other words, a particular embodiment of the invention (the feed being a paraffin-olefin mixture) comprises the following steps (reference is made also to figure 2):

- a) preliminary oligomerization of a (C₄ + C₄ +) mixture at a temperature of from 320 to 380°C using a catalyst bed of zeolitic nature(see e.g. U.S. patent 4,150,062) to obtain a mixture of olefins having from 5 to 8 C atoms, with the C₄ + paraffins remaining unconverted;
- b) cooling and condensation of the oligomerization effluent in order to separate the C₄⁺ paraffins as a gaseous phase, and conversion of the remaining (C₅⁻ -C₈) mixture to propylene under typical "post-pyrolysis" operative conditions;
- c) cooling of the effluent from the reactor for the conversion to propylene and compression of said effluent, preferably at 13-16 absolute Kg/cm², whereby the hydrocarbons having 4 or more C atoms are condensed and the hydrocarbons having less than 4 C atoms are separated as a gaseous phase.

These hydrocarbons ($\langle C_4 \rangle$) can be advantageously recycled to a conventional battery of distillation columns for thermal or catalytic cracking, in order to recover all the propylene contained therein. The small and possible amounts of aromatics (BTX) can be easily separated from the other C_4 hydrocarbons and recycled together with the final $(C_4 - C_8)$ mixture, containing small amounts of butanes produced during the conversion reaction described under item (b) above.

In order to carry out the oligomerization, C₄ olefins, containing C₄ paraffins in any proportion, are initially brought into contact with a catalyst of zeolitic nature, for instance ZSM5 or ZSM11, in an acid or in a m dified form, at 250-400°C (pr ferably 320-380°C) and at space

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velocities from 2 to 10 (preferably 4 - 8) kg/h of reactants per kg of pure zeolite (binder excluded). In other words, the olefins of the $(C_4 + C_4^{-1})$ mixture are converted almost totally into a $(C_5 - C_8^{-1})$ olefinic mixture, while butane and isobutane do not react. The separation of the butanes can thus be carried out very easily by simple cooling with water at room temperature. The olefinic $(C_5 - C_8^{-1})$ mixture liquifies while the butanes are separated as a gaseous phase, said $(C_5 - C_8^{-1})$ mixtures being optimal raw-materials for the production of propylene.

Depending on the operative conditions of the synthesis of the silicalite, as e.g. the dilution of the starting solutions, the resulting crystallites may have a widely variable. Size. The crystallites to be employed in the process according to the invention should have, in general, rather small sizes, such small sizes being obtainable, for instance, following the teachings of U.S. patent 3,926,782. The following examples are given merely for illustration purposes and do not in any way limit the scope of the in vention.

OPERATIVE CONDITIONS COMMON TO ALL EXAMPLES

As to the method of preparation of the catalysts see:

- for silicalite-1: U.S. patent 4,061,724;
- for boralite: Taramasso et al.: "Molecular Sieve Borosilicates", Proc. 5th Int. Conf. on Zeolites, Naples 1980, pages 40-48 (Heyden and Son Ltd. London 1980); the boralite used in the examples has a ratio SiO₂: B₂O₃ of 7
 - (by moles); a more recent method for obtaining boro-silicates is described in European patent publication 77946 .
- for chromosilicates: Italian patent publication 22568 A/82, in the name of the Applicant; the chromosilicate used in the examples has a ratio SiO_2 : Cr_2O_3 of 38 (by moles).

As to zeolites ZSM5 showing a very high SiO₂: Al₂O₃ ratio, not exemplified, see Italian patent publication 21699 A/83, in the name of the Applicant. In the abs nce of different indications, all the catalysts were activated 2 hours at 540°C before being us d.

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EXAMPLES 1-6

3 g of zeolitic catalyst, in admixture with 0.9 g of SiO₂ (as a binder), were loaded into a microreactor which was continuously fed, at a pressure slightly above atmospheric pressure with a 50/50 mixture of butene-2-trans and butene-2-cis. Operative conditions and results are given in Table 1.

0 EXAMPLE 7

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Example 4 was repeated using a silicalite-1 impregnated with a chromium salt, thus obtaining slightly better results.

EXAMPLES 8-15

0.5 g of zeolitic catalyst, in admixture with 0.21 g of Al₂O₃ (as a binder), were loaded into a microreactor which was continuously fed, at a pressure slightly above the atmospheric pressure with a 70/100 mixture of n-pentene and of helium (by moles). Detailed operative conditions and results are given in Table 2. When modified zeolitic catalysts were used, the modifying element was added by means of impregnation.

EXAMPLES 16-23

Examples 8 and 9 were repeated replacing pentene with n-hexene (ex. 16 and 17), with 2-methyl-pentene-1 (ex. 18, 19 and 20) and with n-octene-1 (ex. 21, 22 and 23), respectively, the parameters being slightly modified as indicated in Table 3, which shows also the very good results.

EXAMPLES 24-29

2.12 g of zeolitic catalyst were loaded into a micro-reactor which was continuously fed whith 7.5 kg/h of iso-butene per kg of zeolitic catalyst at 550°C and at a pressure slightly above atmospheric pressure. Data and results are given in Table 4.

EXAMPLES 30 AND 31

Examples 24 and 25 were repeated raising the pressure to 5 absolute atmospheres and the space velocity (WHSV) from 7.5 to 45 h^{-1} , thus obtaining analogous results.

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EXAMPLES 32-35

microreactor which was continuously fed with 60kg/h of the olefins of example 1 per kg of silicalite, at high pressures (6 absolute atmospheres for example 32 and 33 and 8 abs. atm. for ex. 34 and 35). The detailed operative conditions and the results are given in Table 5. The results show clearly that a slight increase in pressure, corresponding approximately to the pressure of the olefins in the industrial manufacturing plants, allows the same conversions and selectivities, but at a much higher velocity. In other words, it is surprising and was quite unexpected that a suitable increase of the pressure increases very much the productivity of the silicate and therefore the output of the industrial plants.

EXAMPLE 36

50 parts by weight of silicalite-1 were admixed with 50 parts of Al₂O₃ (binder) and the mixture was loaded into a microreactor which was continuously fed at atmospheric pressure, with a 50/50 mixture of butene-2-trans and bute-ne-2-cis, at 550°C and at a space velocity of 20 kg/h of olefins per kg of silicalite (binder excluded). The run was very long (120 h) and the results, hour by hour, were continuously monitored and plotted in figure 1. It is worth-while to note that the initial decrease of conversion is reversed after a few hours; thus the conversion of the olefins itself is likely to be a stimulating activation for the catalyst. In other words, catalyst and reaction effect each other by a mutual and beneficial activating action.

EXAMPLE 37

Example 36 was repeated by using a mixture 65% b.w. silicate-1 + 35% b.w. Al_2O_3 and by raising the space velocity to 67 h⁻¹. In this case, the conversion initially decreased and then the phenomenon was reversed.

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5	rea .	60 44 F F F F F F F F F F F F F F F F F F	12.25 40.45 23.65 7.85 30.82	cluded);
10	· и	Chromosilicate {d} sem Ex. 1 i h	29.75 34.07 29.73 19.88 32.25 3.46	Space Velocity) namely Kg/h of olafine per Kg. of pure extalyst (binder excluded); binder; binder; soles)
·	-2	88 88 88 88 88 88 88 88 88 88 88 88 88	77.06 13.47 10.39 45.13 31.75 5.44	a pure sat
15 	r»	Silienlite-1 OO see Ex. 1 " "	61.59 15.26 44.79 49.23 26.15 6,85 4,95	olefine per Kg
7 A B L E	2		29.01 40.40 27.30 18.94 31.73 2.55	nemely Kg/h of
	1	Borelite (b) 9 grams (c) 500 6	45.95 29.79 28.36 28.72 38.76 2.51	Space Velocity) oles); binder; soles)
2 5	EXAMPLE	Cotestive conditions: Catalyst S10_/A1_0 (noles) Amount of catalyst Temp. (oC) Snace velocity (a) Data purvey after:	Results (Rb.m.) Conversion Selectivity to iso-t, Selectivity to C, Yield (iso t, + C, Sel. to compoundaying Sel. to saturated compounds & C, Selectivity to C,	tourly (by m (by m) (b) as 18 (by m)

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	. 15	€.	See Ex. 14	F		£	£	42 90		60.92	19.63	39.71	36.15	A	28.37	1.88	6.05	3.69		[h. 20]201.	(caral (a)		
5	14	· (£)	Cr-Silicalita	8	See Ex. 8	₽.	5	£		69.22	20.51	34.25	37.90	0.41	29.33	1.21	7.79	6.02		7 20 /8 0 - 7	2, 2, 3	• •	
10	13	-	See Ex. 12	=	F	p	Ħ	5 h		10.04	17.92	. 29.52	4.77	0.79	30.38	0.29	4.58	7.27		(binder): (d	F7		
	12		Chromosilicate	*	See Ex. 8	=	= ·	1 h		16.80	21.25	32.50	9.03	59.0	26.90	0.29	7.97	10.83		0.21 0 of A1 0	~	•	
15 Z 3 3 8 Y E	11	·	See Ex. 10	=	E.	#	E	6 h		19.52	27.97	38.93	13.D6	18.0	24.70	0.25	1.79	5.79		Bure zeolita +		gnation.	
20	10		Boralite	Ē	See Ex. 8	F	, #	1 h		42.81	23.43	39.27	75.84	94.0	27.61	0.58	3.59	4.95		: (c) 0.5 a of		Cr. added by means of impregnation.	-
	6		See Ex. 1	=	5	a	r	.		57.00	19.80	39.42	33.76	0.70	29.42	78.0	56.E	3.82		luena + srelanas	•		
25	8		Silicalite-1	8	3	550	7.14	1.5		64,26	19.44	33.19	35,93	98.0	29.27	1.36	6.65	3.06		Benzese + to	38 (by woles).	elning 0.1% b.	•
30	EXAMPLE	Operative Conditions	Catalyst	SIO_/Al_O_(moles)	Amount of catalyst	Temp. (°C)	Space velocity (a)	Data survey after:	Results (% D.u.)	Conversion	Selectivity to C.	Selectivity to C	rield (c2 + c2)	Sel. to estarated	Sel. to C.	Sel. to 1 C.	Sel. to BTX [b]	Sel . to others		(a) See aw 1:(b) RIX = benzene + toluene + zwlenes: (c) 0.5 d of bure zeolite + 0.21 o of Al O (binder): (d) Sin /9 n = 2 (b; selection)	(K) Sio . Cr. 0	(*) Silicalite-1 containing 0.1% b.w.	

		•		,						···		i	D,	-	*			 •	(71	09	U (5 U		-
	•	23	(***)	See Ex. 21		p	909	7.14	. 4		95,50	16.64	27.55	42.23		# ! = :	£.37	21.97	75°C	3.14	n-octene-1				•
5		. 22	(+++)	Sea Lx. 21		E	Þ	, E	.c. 103		97.47	1.60	32.13	37.75			59°0	27.08	2.05	2.77	(4+) Fead = 2-methyl-pantane-1; (49*) Feed = n-ottene-1				
to		21	Î	See Ex. 16	=	E	24	=	4 1		8	9.16	31.64	40.80		30.32	1.06	. 24-14	2.86	3.74	thyl-pantane-1;			٠.	
		0 Z	£	Sae Ex. 18	£	£	±	E	7 h		6.68	7.49	55.35	84*95	. <u>.</u>	1 03	0.00	20.67	0.99	3.26	*) Fead = 2-mei				
15	11.6	61	<u>ŧ</u>	See Ex. 18		F	₽,	F	6.5		97.36	9,29	53.79	61,41	13 6	12.55	0.13	16.54	2.78	3.40	1				
20	1.1	18	£	See Ex. 16	8			=	1 h		97.79	10.55	53,03	52.13	9 6	16.78	0.30	13,82	1.69	2.97	; (*) feed -		-		
		11	€	See Ex. 15	s	F	E	=		·	96.83	9.12	52,90	£0.05	, E	14.80	0.23	17.55	2.22	2.51	luene + Aylenes				
25		91	(*)	Silicalite-1	8	See Ex. 8	550	7.14	4.7		98-01	10.38	51.14	60.2B	9, 0	7 7	14.0	14.64	2,81	2.20	- benzans + to				
30		EXARPLE	Oserative Conditions:	Catalyst	\$10_/A1_0_ (mules)	Amount of catalyst	Temp. (°C)	Space velocity (a)	Data survey efter:	leralts (E h.m.)	Consersion	Selectivity to C_	Selectivity to C.	Yseld (C2 + C3)	Sel. to saturated	4 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 -	Sel. to t.	Sel. to 8TX (6)	Sal.to C	Sel to others	(a) See ex. 1 (b) BIX = benzans + toluene + mylanes ; (*) feed = n-hexane-1;				

										_	.]	11	-			•		01	090	60
																	•	_		
5	29		See Ex. 28	u	=	#	#	5 h		4	12.4	2.2	8,5	(II)		B.1	75.1			
10	82		Chromosilicate	(a)	See Ex. 24	550	7.5	<u>-</u>		in de	15.5	7.4	0.8	5.1		13.7	ē . 60			. .
	7.2		See Ex. 25	=	B	=		£ 9	-	62.4	8-8	15,3	1.2	6.5		4.0	65.7			
15 4.	92		Boralite	(R)	See Ex. 24	250	7.5	#		77.4	39.1	30.3	3.4	5.7		7.8	0.4	(by moles).		
3 1 8 ¥ L	25		See Ex. 24	•	G .	•	F	, <u>.</u>	148 -41	77.3	40.8	31.5	2.6	5.6		7.0	0.77	(b) 510 ₂ : Cr ₂₃ = 38 (by moles).		
	17		Sillcalite-1	ያ	2.12 g	1956	7.5	£		61.8	8.44	35.0	2.4	9-5		4.0	35.1	les); (b) sio		
30	EXAMPLE	Operative conditions:	Latalyst	Sin,/Al,0, (moles)	Asount of catalyst	Tesp (aC)	Space welocity(h])	Usta survey after:	Results (% b.w.)	Conversion	Selectivity to C_	Yield (C_)	Se), to C.	Sel to saturated	carpounds <c< th=""><th>Sel to compounds & C</th><th>Sel to linear betenes</th><th>(a) SiO₂ : B₂O₃ = 7 (by modes);</th><th></th><th>•</th></c<>	Sel to compounds & C	Sel to linear betenes	(a) SiO ₂ : B ₂ O ₃ = 7 (by modes);		•
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5			•																					
10																		,						
	13	£	See Ex. 34	7	=	F	P	H, 8		67.32	28.00	17.50	3.20	8.40		45.60	16.50							
15 	18	(See Ex. 32	•	E	280	99	<u>-</u>		79.31	33.55	31.45	4.23	14.14		27.80	14.19				٠			
20	33	·(*)	Sem Ex. 32	F				-		73.80	39.69	23.44	3.19	10.62		25.73	20.58							
	32	(*)	Sillealite-1	8	69	570	99	=		82.24	31.98	26.30	3.88	12.35	,	36.02	14.76	Atmospheres;	strospheres.					
25 30	ЕХАЙРІЕ	Operative conditions:	Catalyst	S10_/A1_0_ (moles)	Amount of Catalyst	Temp., (aC)	Space velocity (h)	Date sureey after:	Results (% bow.)	Charatsion	Selectivity to C	vield (c_) 3	Sel. to C.	Sel.to saturated compounds	ॐ	Sel. to compounds & C.	Sel. to implutane	(*) Pressure = 6 sbsolute stmospheres;	(**) Pressure = 9 shablute					
																-:-								

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CLAIMS

- 1. A process for the conversion of olefins having from 4 to 12 carbon atoms to propylene, characterized in that said olefins are brought into contact with a zeolitic compound (optionally in admixture with a binder), selected from silicalites, boralites, chromosilicates and those zeolites ZSM5 and ZSM11 in which SiO₂/Al₂O₃ (by moles) is ≥ 350, at a space velocity of from 5 to 200 kg/h of olefins per kg of pure zeolitic compound (binder excluded) and at a temperature of from 400 to 600°C, said zeolitic compound being used as such or in a modified form.
 - 2. The process of claim 1, wherein said compound is silicalite-1, the pressure is substantially atmospheric and the space velocity is from 5 to 50 h^{-1} .
 - 3. The process of claim 1, wherein said compound is silicalite-1, the pressure is from 1.5 to 7.5 absolute atmospheres and the space velocity is from 50 to 200 $\rm h^{-1}$
- 4. The process of any of claims 1-3, wherein said compound is silicalite-1 which is activated in the conversion reaction of said olefins to propylene, under the operative conditions of claim 1, the initial (activating) run being carried out at least 8 and preferably 12 hours.
 - 5. The process of any of claims 1-4, wherein said olefins are selected from isobutene and linear butenes and wherein said binder is SiO₂.
- 55 6. The process of any of claims 1-4, wherein said olefins contain from 5 to 8 carbon atoms and wherein said binder is Al₂O₃.

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- 7. The process of any of claims 1-6, wherein said compound is silicalite-1 in a non-modified form.
- 8. The process of any of claims 1-6, wherein said compound is silicalite-1 in a modified form, the modifying element being Cr, Mg, Ca, Sr or Ba.
- 9. A process for the conversion of olefins having 4 carbon atoms into propylene, characterized in that said olefins are brought into contact with a zeolitic compound (optionally in admixture with a binder) selected from silicalites and those zeolites ZSM5 and ZSM11 in which SiO₂/Al₂O₃ (by moles) is >350, at a pressure of from 1.5 to 7.5 atmospheres, a space velocity of from 50 to 200 kg/h of olefins per kg of pure zeolitic compound (binder excluded) and a temperature of from 400 to 600°C, said compound being used as such or in a form modified by an element selected from Cr, Mg, Ca, Sr and Ba.
 - 10. A process for the conversion of olefins having from 5 to 8 carbon atoms into propylene, characterized in that said olefins are brought into contact with a zeolitic compound (optionally in admixture with a binder) selected from silicalites and those zeolites ZSM5 and ZSMI1 in which SiO₂/Al₂O₃ (by moles) is ≥350, at a pressure of from 1.5 to 7.5 atmospheres, a space velocity of from 50 to 200 kg/h of olefins per kg of pure zeolitic compound (binder excluded) and a temperature of from 400 to 600°C, said compound being used as such or in a form modified by an element selected from Cr, Mg, Ca, Sr and Ba.
- 35 11. The process of claims 9 or 10 wherein said compound is silicalite-1 which is activated in the conversion reaction of said olefins to propylene, under said operative conditions, the initial (activating) run being carried out at least 8 and preferably 12 hours.

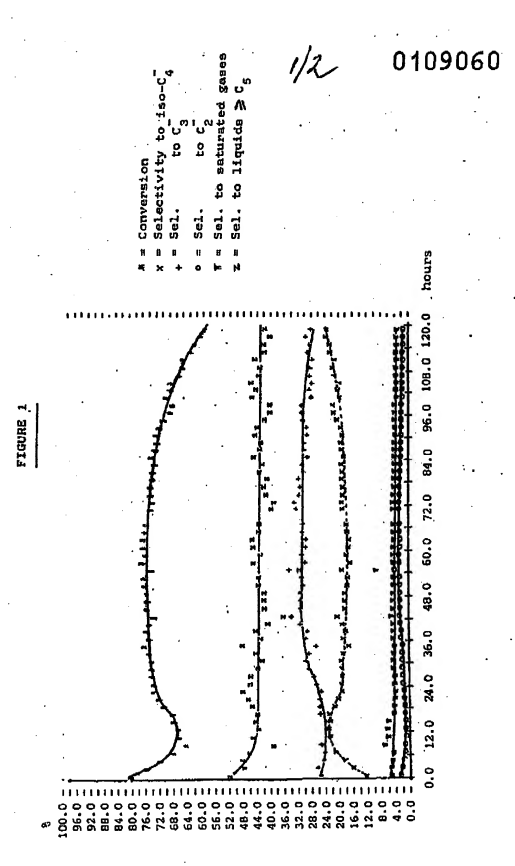
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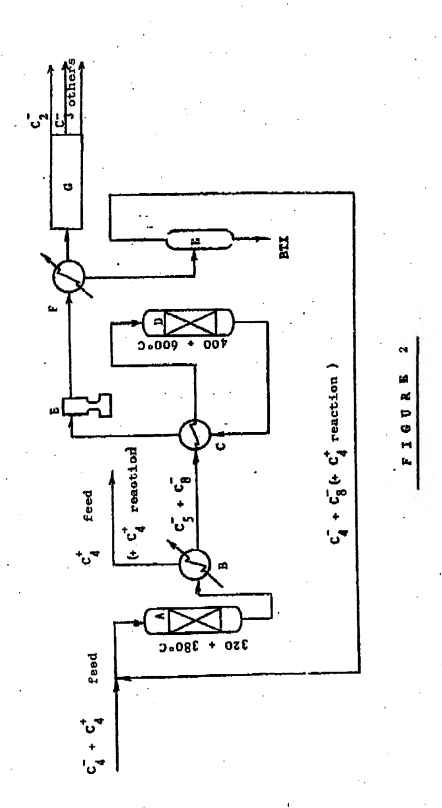
- 12. A process for the conversion into propylene of mixtures of olefins having 4 C atoms, said mixtures containing also ${\bf C_4}^+$ paraffins that cannot be easily separated from olefins, which process comprises the following steps:
 - a) preliminary oligomerization of a $(C_4^- + C_4^+)$ mixture at a temperature of from 320 to 380°C using a catalyst bed of zeolitic nature to obtain a mixture of olefins having from 5 to 8 C atoms, with the C_4^+ paraffins remaining unconverted;
 - b) cooling and condensation of the oligomerization effluent in order to separate the C₄⁺ paraffins as a gaseous phase, and conversion of the remaining (C₅⁻ C₈⁻) mixture to propylene under the typical "post-pyrolysis" reaction conditions of claim 1;
 - c) cooling the effluent from the reactor for the conversion to propylene and compression of said effluent preferably at 13-16 absolute kg/cm², whereby the hydrocarbons having 4 or more C atoms are condensed and the hydrocarbons having less than 4 C atoms are separated as a gaseous phase.

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EUROPEAN SEARCH REPORT

0109060 Application number

EP 83 11 1241

Category	Citation of document wit	BIDERED TO BE RELEVANT th indication, where appropriate, vant passages	Reterant to dalm	CLASSIFICATION OF THE APPLICATION (Int. Ct. 1)
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EUROPEAN PATENT SPECIFICATION

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(5) Int. Cl.⁴: **C 07 C 11/06**, C 07 C 4/10, B 01 J 29/28, B 01 J 29/04

(1) Application number: 83111241.2

② Date of filing: 10.11.83

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(A) Process for the conversion of linear butenes to propylene.

- (3) Priority: 10.11.82 IT 2415382 24.12.82 IT 2497982 14.03.83 IT 2005383 03.05.83 IT 2089683
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Description

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Nowadays huge amounts of ol fin cuts, from C_4^- to C_{12}^- , linear or branched, are available throughout the world and they are widely employed for different purposes such as described, for instance, in Italian patent publications 24152 A/82, 24550 A/82 and 19292 A/83. Sometimes, however, because of contingent reasons, even outside the chemical field such as, for instance, transportation difficulties, it would be better to have still further possibilities of use. A promising use of said olefins would be their conversion into propylene and/or ethylene.

EP—A—37671 discloses a process for the acid catalyzed conversion of larger olefins to smaller olefins, wherein the larger olefins are contacted with a zeolitic compound, e.g. ZSM5 or ZSM11 zeolites, having a silica to alumina mole ratio of at least 12, the zeolitic compound being used as such or in a modified f rm.

However, endurance tests have shown that the excellent initial behavior of some catalysts such as, for instance, ZSM5 and ZSM11, disappears after some time; after a few weeks conversion and selectivity drop to poor levels.

An object of the invention is to provide a process for catalytically converting olefin cuts $C_4 - C_{12}$ with high propylene (and, optionally, also ethylene) yields and with a long life time of the catalyst b for substitution or regeneration.

In its most general form, the invention concerns a process for the conversion of linear butenes to propylene, which comprises contacting said butenes with a zeolitic compound, optionally in admixture with a binder, characterized in that said zeolitic compound is selected from silicalites, boralites, chromosilicates and those zeolites ZSM5 and ZSM11 in which the mole ratio SiO₂/Al₂O₃ is ≥350 and that said conversion reaction is carried out at a temperature of from 500 to 600°C and at a space velocity of from 5 to 200 kg/h-of-butenes per kg of pure zeolitic compound (binder excluded).

The zeolitic compound is used as such or in a modified form.

The behavior of silicalites depends on the conversion pressure; for instance, if the silicalite is silicalite-1 and if the pressure is substantially atmospheric, the space velocity must be lower than $50 h^{-1}$; if said compound is silicalite-1 and if the pressure is from 1.52 to 7.60 bar (1.5 to 7.5 absolute atmospheres) the space velocity must be in general above $50 h^{-1}$.

Best results are obtained when said catalytic silicalite-1 is activated in the conversion reaction of the olefins into propylene, under said operative conditions. This initial (activating) run takes at least 8 and preferably 12 hours; the silicalite-1 is used as such or in a modified form and the modifying element is selected from Cr, Mg, Ca, Sr and Ba.

The modifying element can be incorporated into the catalyst by means of ion exchange or by any other method, for instance impregnation or co-precipitation during the synthesis of the zeolitic compound.

According to a particularly advantageous way for the preparation of a non-modified silicalite, the raw product coming from the zeolite synthesis is dried, for instance at 120°C, calcined (e.g. at 540°C for som hours), in order to remove all the residual organic templating agent, and then exchanged with an aqueous solution of HCl, NH₄Cl, NH₄NO₂ or an equivalent H⁺ or NH₄⁺ source. When an ammonium compound is used, it is necessary to heat, e.g. at 400°C, in order to obtain the acid form of the silicate. A survey of techniques alternative to ion exchange is given e.g. in U.S. patents 3,140,249; 3,140,251; 3,140,253 and in European patent publication 30796, 36707, 37168, 40463, 68754.

The zeolitic material, after calcination and conversion into the acid form, show a long endurance and a very high catalytic activity. These zeolitic compounds can be used as catalysts (as such or in a modified form) optionally in admixture with suitable amounts of binders, for instance SiO₂ or Al₂O₃. A list of oth r binders can be found e.g. in European patent publication 36707. The regeneration can be carried out in air for some hours, at 400—600°C. A steam regeneration is described in European patent publication 36704 and according to a further and very successful method, the catalyst can be regenerated by a hydrogen treatment.

As to the initial activation of the catalyst, some methods are described in European patent publications 34444 and 35830; in general it is advisable to activate the catalyst for some hours in air, at 450—750°C (preferably 540—700°C). Furthermore the conversion itself (of olefins to propylene) has an activating effect on the zeolitic catalyst. In other words, catalyst and reaction affect each other in a mutual, reciprocal and beneficial activating reaction.

Any process for the conversion of more or less heavy olefinic cuts into propylene will be indicated hereafter, as a "post-pyrolysis" process. When the feed of a post-pyrolysis process is a mixture of olefins having 4 C atoms, there is a considerable problem to be solved, because the C₄ cuts always contain substantial amounts of paraffins, in general, also having 4 C atoms, which paraffins pass (at 400—600°C) the zeolitic bed without taking part in any reaction. Furthermore a small amount of C₄ paraffins is produced by the post-pyrolysis process itself. The conversion to C₃H₆ could be enhanced by a recycle of the non-reacted C₄ olefins or of the C₄ olefins formed during the reaction. In such a case, howev r, an increasing accumulation of n-butan and of isobutane would tak place. This drawback could be avoided by a separation of paraffins from ol fins b fore feeding the reactor but such a separation is rather difficult. But nes and isobutane cannot be is lated by a simple distillation and it is usually necessary t carry ut an extractive distillation (a complicated technique), which is particularly burdensome for the C₄ cuts coming from catalytic cracking, where butan and isobutane may account for even 50% of the whole. The problem,

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however, can be solved in a surprisingly easy way by the process according to the invention, whon employing an integrated oligomerization step. In other words, a particular embodiment of the invention (the feed being a paraffin-olefin mixture) comprises the following steps (reference is made also to figur 2):

a) preliminary oligomerizati n of a $(C_4^- + C_4^-)$ mixtur at a temperature of fr m 320 t 380°C using a catalyst b d of zeolitic natur (see .g. U.S. pat nt 4,150,062) to btain a mixtur of olefins having from 5 to 8 C atoms, with the C_4^+ paraffins remaining unconverted;

b) cooling and condensation of the oligomerization effluent in order to separate the C_4^+ paraffins as a gaseous phase, and conversion of the remaining $(C_5^--C_8^-)$ mixture to propylene under typical "post-pyrolysis" operative conditions;

c) cooling of the effluent from the reactor for the conversion to propylene and compression of said effluent, preferably at 13—16 bar, whereby the hydrocarbons having 4 or more C atoms are condensed and

the hydrocarbons having less than 4 C atoms are separated as a gaseous phase.

These hydrocarbons (<C₄) can be advantageously recycled to a conventional battery of distillation columns for thermal or catalytic cracking, in order to recover all the propylene contained therein. The small and possible amounts of aromatics (BTX) can be easily separated from the other C₄ hydrocarbons and recycled together with the final (C₄ -C₈) mixture, containing small amounts of butanes produced during the conversion reaction described under item (b) above.

In order to carry out the oligomerization, C_4 olefins, containing C_4 paraffins in any proportion, are initially brought into contact with a catalyst of zeolitic nature, for instance ZSM5 or ZSM11, in an acid or in a modified form, at 250—400°C (preferably 320—380°C) and at space velocities from 2 to 10 (preferably 4—8) kg/h of reactants per kg of pure zeolite (binder excluded). In other words, the olefins of the $(C_4^-+C_4^-)$ with with the properties of the carried out very easily by simple cooling with water at room temperature. The olefinic $(C_5^--C_8^-)$ mixture liquifies while the butanes are separated as a gaseous phase, said $(C_5^--C_8^-)$ mixtures being optimal raw-materials for the production of propylene.

Depending on the operative conditions of the synthesis of the silicalite, as e.g. the dilution of the starting solutions, the resulting crystallites may have a widely variable size. The crystallites to be employed in the process according to the invention should have, in general, rather small sizes, such small sizes being obtainable, for instance, following the teachings of U.S. patent 3,926,782. The following examples are given

merely for illustration purposes and do not in any way limit the scope of the invention.

Operative Conditions Common to all Examples

As to the method of preparation of the catalysts see:

- for silicalite-1: U.S. patent 4,061,724;

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— for boralite: Taramasso et al.: "Molecular Sieve Borosilicates", Proc. 5th Int. Conf. on Zeolites, Naples 1980, pages 40—48 (Heyden and Son Ltd. London 1980); the boralite used in the examples has a ratio SiO₂:B₂O₃ of 7 (by moles); a more recent method for obtaining boro-silicates is described in European patent publication 77946;

— for chromosilicates: Italian patent publication 22568 A/82, in the name of the Applicant; the

40 chromosilicate used in the examples has a ratio SiO₂:Cr₂O₃ of 38 (by moles).

As to zeolites ZSM5 showing a very high SiO₂:Al₂O₃ ratio, not exemplified, see Italian patent publication 21699 A/83, in the name of the Applicant. In the absence of different indications, all the catalysts were activated 2 hours at 540°C before being used.

Examples 1—6

3 g of zeolitic catalyst, in admixture with 0.9 g of SiO₂ (as a binder), were loaded into a microreactor which was continuously fed, at a pressure slightly above atmospheric pressure with a 50/50 mixture of butene-2-trans and butene-2-cis. Operative conditions and results are given in Table 1.

Example 7

Example 4 was repeated using a silicalite-1 impregnated with a chromium salt, thus obtaining slightly better results.

Examples 8-11

3 g of silicalite-1, without binder, were loaded into a microreactor which was continuously fed with 60 kg/h of the olefins of example 1 per kg of silicalite, at high pressures (6.08 bar for example 8 and 9 and 8.1 bar for example 10 and 11). The detailed operative conditions and the results are given in Table 2. The results show clearly that a slight increase in pressure, corresponding approximately to the pressure of the olefins in the industrial manufacturing plants, allows the same conversions and selectiveness, but at a much higher velocity. In other words, it is surprising and was quite unexpected that a suitable increase of the pressure increases very much the productivity of the silicate and therefore the output of the industrial plants.

Example 12

50 parts by weight of silicalit -1 w re admixed with 50 parts of Al₂O₃ (binder) and the mixture was

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loaded into a microreactor which was continuously fed at atmospheric pressure, with a 50/50 mixture of butene-2-trans and butene-2-cis, at 550°C and at a space velocity of 20 kg/h of olefins per kg of silicalite (binder excluded). The run was very long (120 h) and the results, hour by hour, were continuously monitored and plotted in figure 1. It is worthwhile to note that the initial decrease of conversion is reversed after a few hours; thus the conversion of the ol fins itself is likely to b a stimulating activation for the catalyst. In other words, catalyst and reaction effect each other by a mutual and beneficial activating action.

Example 13

Example 12 was repeated by using a mixture 65% b.w. silicate-1+35% b.w. Al₂O₃ and by raising th space velocity to 67 h⁻¹. In this case, the conversion initially decreased and then the phenomenon was reversed.

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	9		See Ex. 5	:	:	:		u 9		12.25	40.46	23.65	7.85	30.82	4.77	0.31
	ည		Chromosilicate	(p)	See 3x. 1	:	:	-		29.75	34.07	29.73	19.88	32,29	3.46	0.45
	а-маст. 4		See Ex. 3	8		60 60		4 2		77.06	19.47	40.39	45.13	31.75	5.44	2.95
E 1	8		Silicalite-1	8	See Ex. 1			د		81.99	15.26	44.79	49.23	28.15	6.85	4.95
TABLE 1	2		See Ex. 1	:	=		:	5 h		29.01	40.40	24.90	18.94	31.73	2,55	0.43
	-		Boralite	(q)	3 grams (c)	200	9	£		45.95	29.79	28.36	26.72	38.76	2.51	0.57
	Example	Operative conditions:	Catalyst	SiO ₂ /Al ₂ O ₃ (moles)	Amount of catalyst	Temp. (°C)	Space velocity (a)	Data survey after:	Results (% b.w.)	Conversion	Selectivity to iso-C	Selectivity to C3	Yield (iso $C_4^- + C_3^-$)	Sel. to compounds \gtrsim $C_{\rm S}$	Sel. to saturated compounds < C	Selectivity to C_2

(a) WHSV (Weight Hourly Space Velocity) namely Kg /h of olefine per Kg. of pure catalyst (binder excluded);

⁽b) $SiO_2 : B_2O_3 = 7$ (by moles); (c) + 0.9 g of SiO_2 as binder; (d) $SiO_2 : Cr_2O_3 = 38$ (by moles).

TABLE 2

#	(**)	See Ex. 10	:	:	:	:	8 h		67.32	26.00	17.50	3.20	8.40		45.60	16.50
10	(**)	See Ex. 8	•	•	280	09	- م		79.31	39.66	31.45	4.21	14.14		27.80	14.19
6	(.)	See Ex. 8		Table over 100	 2	:	8 1		73.80	39.89	29.44	3,19	10.62		25.73	20.58
8	(.)	Silicalite-1	8	8	920	09	-		82.24	31.98	26.30	3.89	13.35		36.02	14.76
Example	Operative conditions:	Catalyst	SiO ₂ /AI ₂ O ₃ (moles)	Amount of catalyst	Temp. (°C)	Space velocity (h ⁻¹)	Data survey after:	Results (% b.w.)	Conversion	Selectivity to C ₃	Yield (C3)	Sel. to C ₂	Sel. to saturated compounds	*C4	Sel. to compounds ≽C ₅	Sel. to isobutene

(*) Pressure = 6 absolute atmospheres (6.08 bar).

(**) Pressure = 9 absolute atmospheres (9.12 bar).

Claims

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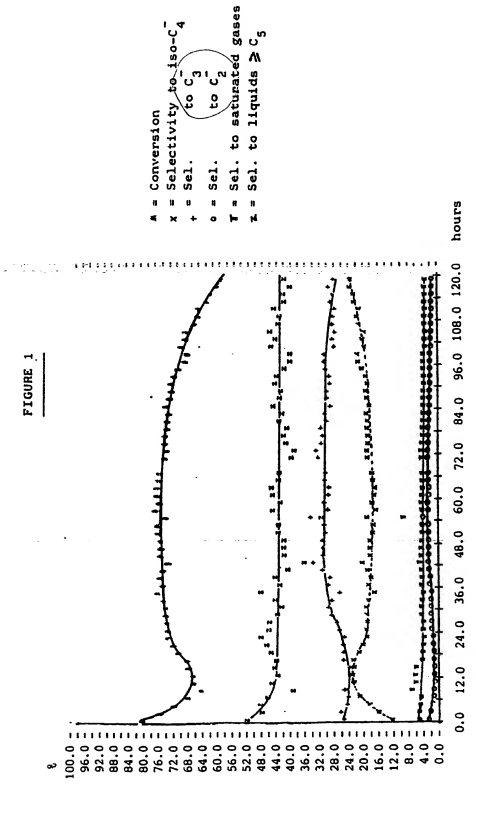
- 1. A process for the conversion of linear butenes to propylene, which comprises contacting said butenes with a zeolitic compound, optionally in admixture with a binder, characterized in that said zeolitic compound is selected from silicalites, boralites, chromosilicates and those zeolites ZSM5 and ZSM11 in which the mole ratio SiO₂/Al₂O₃ is ≥350 and that said conversion reaction is carried ut at a temperature of from 500 to 600°C and at a space velocity of from 5 to 200 kg/h of butenes per kg of pure zeolitic compound (binder excluded).
 - 2. The process of claim 1 wherein said zeolitic compound is silicalite-1.
- 3. The process of claim 2 wherein the space velocity is from 5 to 50 h^{-1} , when the reaction pressure is substantially atmospheric, and is from 50 to 200 h^{-1} , when the reaction pressure is from 1.5 to 7.5 bar.
- 4. The process of claim 2 or 3 wherein the silicalite-1 is activated by the conversion reaction of said butenes to propylene under the reaction conditions of claim 1, the initial (activating) run being carried out at least 8 and, preferably, 12 hours.
 - 5. The process of any one of claims 2 to 4 wherein said silicalite-1 is in a non-modified form.
- 6. The process of any one of claims 2 to 4 wherein said silicalite-1 is in a modified form, the modifying element being selected from Cr, Mg, Ca, Sr or Ba.
 - 7. The process of any one of claims 1 to 6 wherein said binder is SiO₂.

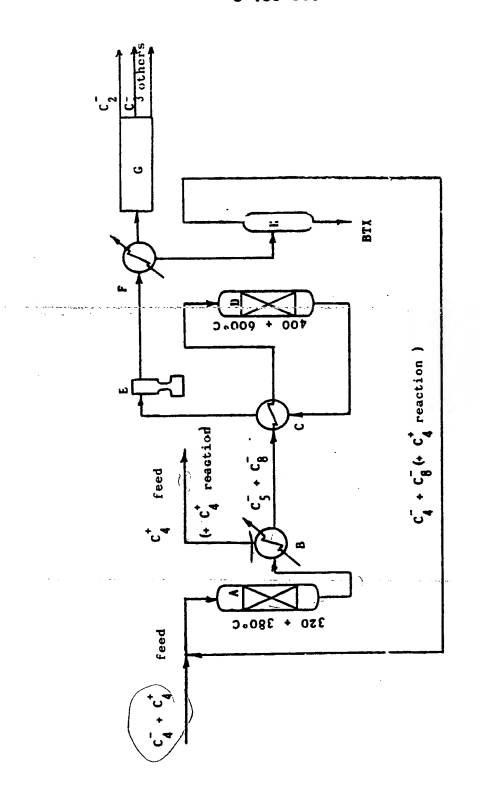
Patentansprüche

- 1. Verfahren zur Umwandlung von linearen Butenen in Propylen, bei dem die Butene mit einer zeolithischen Verbindung, gegebenenfalls im Gemisch mit einem Bindemittel, in Kontakt gebracht werden, dadurch gekennzeichnet, daß die zeolithische Verbindung ausgewählt ist unter Silicaliten, Boraliten, Chromosilicaten und Zeolithen ZSM5 und ZSM11, bei denen das Molverhältnis SiO₂/Al₂O₃ ≥350 ist, und daß die Umwandlungsreaktion bei einer Temperatur von 500 bis 600 °C und einer Raumgeschwindigkeit von 5 bis 200 kg/h Butenen pro kg reiner Zeolithischer Verbindung (Bindemittel ausgenommen) durchgeführt wird.
 - 2. Verfahren nach Anspruch 1, worin die zeolithische Verbindung Silicalit-1 ist.
- 3. Verfahren nach Anspruch 2, worin die Raumgeschwindigkeit 5 bis 50 h⁻¹ beträgt, wenn der Reaktionsdruck im wesentlichen Atmosphärendruck ist, und 50 bis 200 h⁻¹ beträgt, wenn der Reaktionsdruck 1,5 bis 7,5 bar ist.
- 4. Verfahren nach Anspruch 2 oder 3, worin der Silicalit-1 durch die Umwandlungsreaktion der Butene im Propylene unter den Reaktionsbedingungen von Anspruch 1 aktiviert wird, wobei der anfängliche (aktivierende) Durchgang mindestens 8 und vorzugsweise 12 Stunden durchgeführt wird.
- 5. Verfahren nach irgendeinem der Ansprüche 2 bis 4, worin der Silicalit-1 in nicht-modifizierter Form vorliegt.
- 6. Verfahren nach irgendeinem der Ansprüche 2 bis 4, worin der Silicalit-1 in modifizierter form vorliegt, wobei das modifizierende Element ausgewählt ist unter Cr, Mg, Ca, Sr oder Ba.
 - 7. Verfahren nach irgendeinem der Ansprüche 1 bis 6, worin das Bindemittel SiO2 ist.

Revendications

- 1. Un procédé de conversion de butènes linéaires en propylène, qui comprend le contact des butènes avec un composé zéolitique, éventuellement associé à un liant, caractérisé en ce que le dit composé zéolitique est choisi parmi les silicalites, les boralites, les chromosilicates et les zéolites ZSM5 et ZSM11 dans lesquelles le rapport molaire SiO₂/Al₂O₃ est supérieur ou égal à 350 et en ce que la dite réaction de conversion est mise en oeuvre à une température comprise entre 500°C et 600°C et à une vitesse spatiale comprise entre 5 et 200 kg/h de butènes par kilogramme de composé zéolitique pur (liant exclus).
 - 2. Le procédé selon la revendication 1 dans lequel ledit composé zéolitique est la silicalite-1.
- 3. Le procédé selon la revendication 2 dans lequel la vitesse spatiale est comprise entre 5 et 50 h⁻¹, lorsque la pression réactionnelle est de l'ordre de la pression atmosphérique et est comprise entre 50 et 200 h⁻¹, lorsque la pression réactionnelle est comprise entre 1,5 et 7,5 bars.
- 4. Le procédé selon la revendication 2 ou 3 dans lequel, dans les conditions de réaction de la revendication 1, la silicalite-1 est activée par la réaction de conversion des dits butènes en propylène, la procédure initiale (activation) étant effectuée en au moins 8 heures, de préférence 12 heures.
 - 5. Le procédé selon la revendication 1 dans lequel la silicalite-1 est sous une forme non modifiée.
- 6. Le procédé selon une quelconque des revendications 2 à 4 dans lequel ledit silicalite-1 est sous une forme modifiée, l'élément modificateur consistant en Cr, Mg, Ca, Sr ou Ba.
 - 7. Le procédé s l n une quelconqu des revendications 1 à 6 dans lequel ledit liant est formé de SiO₂.





FIGURE